11) Publication number:

0 308 143 A1

(12)

EUROPEAN PATENT APPLICATION

21 Application number: 88308335.4

(1) Int. Cl.4 C03C 25/02 , G02B 6/44

2 Date of filing: 09.09.88

Priority: 18.09.87 US 98253

Date of publication of application:
 22.03.89 Bulletin 89/12

Designated Contracting States:
DE FR GB NL

7) Applicant: AMERICAN TELEPHONE AND TELEGRAPH COMPANY 550 Madison Avenue New York, NY 10022(US)

② Inventor: DiMarcello, Frank Vincent RD2, Annandale
New Jersey 08801(US)
Inventor: Huff, Richard Garner
83 Highland Avenue
Basking Ridge New Jersey 07920(US)
Inventor: Lemaire, Paul Joseph
18 Ferndale Road
Madison New Jersey 07940(US)
Inventor: Walker, Kenneth Lee
1003 Central Avenue
New Providence New Jersey 07974(US)

Representative: Johnston, Kenneth Graham et al AT&T (UK) LTD. AT&T Intellectual Property Division 5 Mornington Road Woodford Green Essex, IG8 OTU(GB)

(54) Hermetically sealed optical fibers.

⑤ An hermetically coated optical fiber is produced by contacting a hot fiber with an organic material such as acetylene. The heat of the fiber causes decomposition and results in an hermetic, carbonaceous coating. This coating is essentially impermedable to both water and hydrogen.

EP 0 308 143 /

HERMETICALLY SEALED OPTICAL FIBERS

Background of the invention

Technical Field

This invention relates to optical fiber and, in particular, to coated optical fiber.

Art Background

Typically, an optical fiber, after it is drawn from a preform, is coated with at least one, and typically two, polymer coatings. These coatings are applied by directing the fiber through a reservoir containing a suitable monomer, drawing the coated fiber through a die, and then curing the monomer into a polymer through exposure to radiation, e.g., ultraviolet radiation. The resulting coatings significantly enhance the mechanical and optical properties of the fiber.

Despite the advantages of polymeric coatings for optical fiber, they are generally permeable to water and hydrogen. This permeation by environmental water or by hydrogen generated during reactions of cable components in applications such as oil well logging or undersea systems has been found to have significant effects. In particular, the interaction of water with the surface of the silica fiber produces surface modifications that lower the fracture resistance of the fiber to applied stress. The Interaction of the silica fiber with hydrogen produces an attenuation in the signal carried by the fiber. Thus, the reliability of the optical fiber, especially in adverse environments or the suitability of the fiber for applications where signal attenuation is not acceptable, necessitates a fiber with a hermetic coating alone or in combination with the typical polymer coatings.

Despite this desire, the deposition of hermetic coatings without substantial degradation of properties or substantial increase in cost is extremely difficult to attain. Various attempts have been made to achieve an economic, hermetic coating. For example, in one approach after draw the fiber is directed through a furnace containing a gas. The furnace induces pyrolytic decomposition of the gas which, in turn, produces a coating on the fiber. The composition of the fiber coating depends on the gas employed. Attempts have been made using organic gases to make hermetic coatings that are primarily carbon compositions. In all these attempts the coating was non-adherent and/or non-hermetic. Such attempts have been described in U.S. Patent

4,512,629 issued April 23, 1985, where C₄H₁₀ gas was employed; in the SPIE Proceedings on Reliability Considerations in Fiber Optic Applications, September 25-26, 1986, Cambridge, Massachusetts, p. 27, where a C₄H₁₀ gas was employed; in the Proceedings of the Optical Fiber Conference, Phoenix, Arizona, 1982, paper WCC1, where the gas utilized was not disclosed; and in Physics of Fiber Optics, Advances in Ceramics, eds. B. Bendow and S. S. Mitra, Vol. 2, pp. 124-133, American Ceramics Society, 1981, where an ion plasma deposition was utilized.

Compositions other than carbon have been produced using the previously described furnace approach for producing hermetic coatings by a gas phase reaction. For example, a combination of C4H10 and TiCl4 has been utilized to obtain titanium carbide coatings while a combination of silane and ammonia has been utilized to obtain silicon oxynitride coatings. (See SPIE Proceedings on Reliability Considerations in Fiber Optic Applications, September 25-26, 1986, Cambridge, Massachusetts, p. 27, and U. S. Patent No. 4,512,629, respectively.) The resistance to static fatigue of these coatings, i.e., on the order of n = 30 - 100, for some applications is not entirely acceptable. Additionally, these coatings have typically been applied at draw rates slower than approximately 1 meter per second. Thus, even if the properties of these non-carbonaceous coatings are acceptable, the relatively slow draw speeds compared to typical speeds of 4-6 m/sec substantially increases cost.

Another approach suggested for producing a non-permeable coating employs the heat associated with the fiber after it is drawn to induce decomposition of a gas and subsequent fiber coating. (This approach is mentioned in U. S. Patent No. 4,575,463 but specifics such as deposition conditions or useful coating precursor gases are not discussed.) Thus, although hermetically coated optical fibers for many applications are desirable, achieving acceptable, economic results is extremely difficult.

Summary of the Invention

An excellent hermetic coating for an optical fiber is attained by inducing decomposition of a suitable organic gas at the fiber surface. For example, if the fiber directly after draw is treated with acetylene, decomposition at the fiber surface occurs, and a carbonaceous coating is produced that is strongly adherent, that does not substantially

degrade optical properties, and which is essentially impermeable at room temperature to water and hydrogen.

Use of a combination of acetylene and a chlorine-containing gas such as chlorine or trichloroethylene has produced even better results. The initial fiber after coating has a loss of 0.38 dB/km at 1.3 μ m and subjecting this fiber to hydrogen at elevated temperatures does not substantially increase this loss. The introduction of chlorine, it is believed, scavenges any hydrogen present during the coating procedure and therefore reduces losses associated with trapped hydrogen. A variety of organic materials produces the desired carbonaceous coating provided decomposition is induced at the fiber surface.

Additionally, it has been found that there is a significant amount of hydrogen containing gas trapped under the hermetic coating upon coating. For relatively non-permeable coatings it is possible that this gas has significant consequences.

Brief Description of the Drawings

FIGS. 1-3 are illustrative of apparatuses suitable for the practice of the invention.

Detailed Description

The procedure for drawing the fiber, and subsequently coating, if desired, with polymer coatings, is performable by conventional techniques. For example, the procedure described in F. V. DiMarcello, C. R. Kurkjian, and J. C. Williams, "Fiber Drawing and Strength Properties" in Optical Fiber Communications, Vol. 1, T. Li, ed., Academic Press Inc., 1985, is utilized to draw the fiber from a preform, and procedures such as described at in F. V. DiMarcello, C. R. Kurkjian, and J. C. Williams, supra, if desired, are utilized to coat the fiber with one or more polymer coatings. Suitable polymeric coatings are extensively discussed in L. L. Blyler, Jr., "Polymer Coatings for Optical Fibers Used in Telecommunications," Polymer News, Vol. 8, 1981, pp. 6-10.

The hermetic coating is formed after drawing the fiber from the preform but before organic polymer coatings are applied. The hermetic coating should interact directly with the surface of the glass fiber to produce the desired adhesion required for hermeticity. Thus, it is not appropriate to deposit the typical polymeric coating(s) before formation of the hermetic material.

The hermetic coatings involved in the Inventive procedure are formed by the interaction of an organic material with the surface of the hot fiber. The fiber should be sufficiently hot to Induce bonding between a carbon atom of the organic gas and a silicon atom of the glass fiber. Although the fiber temperature range required to achieve this result varies depending on 1) the organic gas being employed, 2) the heat of reaction of this gas with the silicon composition of the fiber and, 3) the specific thermal conditions associated with the fiber, typically temperatures in the range 700°C to 900°C are suitable.

The thermal conditions and the quality of the resulting coating are primarily influenced by the rate of fiber draw and the distance from the draw furnace at which the organic gas is interacted with the fiber surface. For example, if acetylene is utilized as the organic composition and interacted with the fiber at a distance of approximately 10 lnches from the hot zone of a draw furnace having a temperature of 2300°C, draw rates below 3.5 meters per second do not produce an adherent coating. Coatings formed under the same conditions but with draw speeds between 3.5 and 6.5 meters per second produce excellent hermeticity.

The thermal conditions not only strongly affect the bonding of organic gas moieties to silicon in the glass fiber but also affect crosslinking between these bonded moieties. If no unsaturated bonds are present in the crosslinked network, a diamond-like structure is formed that is too brittle to afford acceptable mechanical properties for the final fiber and to open sufficiently to allow significant diffusion of water and hydrogen. The lack of unsaturated bonds is typically produced by an excessive fiber temperature. In the previous example involving acetylene, if draw speeds above 6.5 meters per second are utilized, an unacceptable diamond-like film is produced. (A diamond-like film is categorized by being optically transparent, electrically insulating, and primarily having only single bonded carbon as shown by Raman microprobe.) Parameters such as draw speed and contact point of the gas with the fiber are interrelated. The further the contact point from the draw furnace and/or the slower the draw speed the less the heat available for reaction. A control sample is utilized to determine a suitable contact point for a desired draw speed to obtain a hermetic, non-diamond coating.

Although a variety of organic materials lead to advantageous coatings, the level of permeability to hydrogen and/or water varies somewhat. Alkynyls, e.g., lower alkynyls such as acetylene, appear to yield the lowest degree of permeability. It is believed that this result occurs because the reaction of triple bonds upon pyrolysis yields a crosslinked network having unsaturated bonds and because this crosslinking reaction is exothermic. The heat liberated upon breaking of the triple bond contributes to the bonding of the carben of the acetylene

to the silicon of the glass.

Nevertheless, alkyls, e.g., lower alkyls such as propane, and alkenyls, e.g., lower alkenyls such as butadiene, also yield materials that although more permeable to water and hydrogen still have a lack of permeability that is acceptable for many applications. The degree of hermeticity depends on the gas employed. Both the gas to which the coating is to be hermetic and the degree of this hermeticity are important. In applications for which attenuation is most significant permeation to hydrogen is the primary consideration. For these applications, it is desirable to maintain a hermeticity such that the loss measured during continuous exposure to 1 atm. of hydrogen at 1.24 micrometers and at the temperature of use increases less than 0.05 dB/kilometers over an exposure time period that is 1/140 of the desired fiber lifetime. For more demanding applications the loss should not increase more than 0.05 dB/km in an exposure time period shorter than 1/45 of the desired lifetime, and for the most demanding applications such as undersea telecommunications the loss should not increase more than 0.05 dB/km in an exposure time period shorter than 1/20 of the desired lifetime. The factors of 1/140, 1/45 and 1/20 correspond to a hydrogen permeation at the end of the fiber lifetime of 50 percent of saturation 20 percent, and 10 percent respectively (assuming a slow reaction of hydrogen with the fiber compared to the rate of diffusion of hydrogen through the coating.) In situations where a getterer is present in the fiber, greater permeation to hydrogen is acceptable. Indeed, for the most reactive getterers at low hydrogen concentration, e.g., 10⁻⁵ atm., experienced by fibers within some terrestrial cables a hermetic coating is not essential. However, even for the most reactive getterers at higher concentrationsabove 10⁻³ atm. such as in undersea cable and above 0.5 atm. such as experienced in some applications involving cables susceptible to galvanic corrosion--it is desirable to employ a coating which in the absence of the getterer by the measurement described above undergoes a 0.05 dB/km increase no faster than 1/2000 and 1/140 the desired life respectively.

In applications where resistance to static fatigue is most significant, e.g., fibers to be used in severe mechanical conditions, permeation to water or OH radicals is the primary concern. In these applications the static fatigue stress corrosion susceptibility factor should be greater than 70, preferably greater than 150 and the tensile strength should be greater than 400,000 preferably greater than 500,000. It is also desirable for some applications that the combined criteria for static fatigue, tensile strength, and loss all be satisfied.

Irrespective of the organic material employed

and the ultimate use, the object is to contact the fiber at a temperature that produces chemical bonding between carbon and silicon atoms and that produces a crosslinked carbon network having unsaturated bonds. (It is possible to introduce some entities other than carbon atoms into this network. However, generally such atoms decrease hermeticity relative to hydrogen permeation. Thus, such non-carbon entities should be limited to a degree that unacceptable loss is not produced.)

6

It has been found that hydrogen containing entities are present under the hermetic coating after fiber drawing. The exact reason for this phenomenon has not been precisely determined. Possibly because the organic gas employed typically has hydrogen atoms, there is a tendency to trap some hydrogen gas under the hermetic coating/glass fiber boundary. Alternatively, reaction with hydrogen evolved from the glass fiber could be the cause. When the coating is hermetic or presents a barrier to diffusion of this trapped gas (i.e., the diffusion coefficient is greater than (K) (2x10⁻¹²cm²/sec)) for a 1000 Angstrom thick coating at 250 degrees C, where K is the solubility of hydrogen in the coating divided by the solubility of hydrogen in silica, it is possible for it to have a sufficient residence time for substantial reaction with the fiber and degradation of fiber properties. That is, there is sufficient trapped hydrogen and the reaction rate with the fiber at the ambient temperature is sufficiently high relative to its outdiffusion rate through the fiber coating to produce undesirable losses.

Although for many applications the degradation due to trapped gas is acceptable, for more demanding applications such as undersea communication systems, it is desirable to prevent losses greater than 0.03 dB/km from this source. To achieve this goal, the level of trapped gas is reduced or it is removed before substantial reaction with the fibers. The reduction of trapped hydrogen or water is accomplished, for example, by introducing a gettering substance (e.g., chlorine, bromine and/or fluorine), for entities such as hydrogen atoms in the precursor gas. This introduction is advantageously accomplished by using a supplementary gas having chlorine, bromine and/or fluorine atoms. For example, in the use of acetylene, the acetylene is mixed with a gas such as trichloroethylene or molecular chlorine. The level of trapped hydrogen and water through the presence of chlorine is substantially reduced.

At high chlorine levels, e.g., above 1:1 molar ratio of chlorine to acetylene or 1:10 molar ratio of trichloroethylene:acetylene, some chlorine is introduced into the hermetic coating and mechanical properties are to an extent degraded. The chlorine is advantageously removed by heating the fiber to

8

temperatures in the range 900 °C to 1100 °C after hermetic coating but before further coating. Alternatively, the fiber is sequentially treated for a relatively short distance with 1) a mixture of a getter containing gas with organic gas, e.g., 2:1 trichloroethylene:acetylene and then 2) with organic gas, e.g., acetylene. The first gas combination introduces the getterer close to the fiber and the second augments the coating without incorporating excess chlorine.

Another procedure for removing trapped hydrogen containing gas involves introducing a getterer in the fiber at a spatial location that does not degrade optical properties but allows reaction with the trapped gas. Generally, the introduction of dopants such as AI, Ge, and/or P or defects such as draw induced defects in the region that is a distance of more than 10 µm from the center of the core in a fiber having a cladding diameter of 125 µm and a core diameter of 8 µm allows reaction of the dopant or defect with the trapped gas but does not degrade optical properties.

Although the temperature associated with the fiber is sufficient to induce the desired reaction to form the coating, the use of an external furnace to supply some heat is not precluded. Nevertheless, the level of this supplemental heat should not be sufficient to prevent reaction between carbon atoms in the organic gas and silicon in glass fiber composition. If this surface reaction is prevented by excessive inducement of gas phase reaction, sufficient adherence and thus sufficient hermeticity is precluded. Additionally, even if bonding occurs, the particle formation associated with gas phase reactions tend to produce voids in the hermetic coating and thus tends to degrade its properties.

The concentration of the organic material in the vicinity of the fiber also affects the quality of the hermetic coating. Typically, chemical concentrations of 5-10 mole percent in a flow of 3 /min N2 yield desirable results. If the chemical concentration becomes too high it is possible to produce a gas phase reaction, obtain an explosive mixture, and/or generate an excessive concentration of particles, while if the chemical concentration is too low hermetic coating is not achieved. The size of the vessel employed to introduce the gas affects the results by affecting the flow pattern. For vessels greater than 1 inch in diameter, the flow pattern is such that the effective concentration at the fiber is lowered. Therefore, it is advantageous to use a vessel less than 1 inch in diameter. A controlled sample is easily employed to determine an appropriate flow rate for the conditions utilized.

The following examples are illustrative of conditions suitable for the practice of the inventive technique.

Example 1

A draw tower configuration as schematically illustrated in FIG. 1 was employed. This draw tower included a preform feed mechanism, 10, a draw furnace, 11, a monitor for measuring fiber diameter, 13, a coating chamber, 15, a polymer coating dle, 17, a curing station, 19, a coating diameter monitor, 4, and a capstan and takeup mechanism, 16. These components were conventional and have been described in F. V. DiMarcello, supra. A single mode collapsed fiber preform, 12, was inserted in the preform furnace. The furnace was heated to a temperature of approximately 2300 degress Centigrade. Inlet and outlet purges of nitrogen were established by introducing a nitrogen gas flow of 2 /min at inlet, 21 and 22, shown in the enlarged view (FIG.2) of the reaction chamber. Additionally, the exhaust, 25, was initiated and a nitrogen flow through reactant inlet, 26, at a rate of 3.5 /min was introduced. (The inlet and outlet gas purges prevented atmosphere oxygen from entering the reaction chamber and to an extent stripped the boundary layer accompanying the fiber.) The coating applicator for the final polymeric coating was filled with a conventional acrylate UV curable coating. The fiber was initially drawn and threaded through fiber diameter monitor, 13. The fiber was then threaded through stripper plates, 24 and 23, each having an opening of 0.100 inches with a spacing between the stripper plates of 0.25 inches. Fiber threading was continued through 1) the purge outlet, 2) the polymeric coating die, 17, of the coating applicator, 3) the curing station which had two UV lamps providing a power of approximately 300 watts/inch over a length of approximately 18 inches, and 4) the coating diameter monitor to the capstan and takeup mechanism.

The reaction chamber was positioned approximately 10 inches from the hot zone of the furnace and had a cylindrical reaction region measuring approximately 9 Inches long with a diameter of approximately 0.4 inches. A composition containing acetylene, molecular chlorine, and molecular nitrogen having respective flow rates of 185 cc/min, 125 cc/min, and 3.5 /min (measured utilizing an Applied Materials mass flow controller) was introduced into port, 26, as the fiber draw speed was increased to a rate of approximately 5 meters per second. (The fiber temperature as it entered the reaction chamber as measured with an optical pyrometer was approximately 880 degrees C.)

Approximately 9000 meters of fiber were drawn. A series of 10 centimeter lengths from this fiber was measured utilizing a tensile tester made by Instron Corporation. The average measured strength was approximately 600,000 psi. The stress corrosion susceptibility factor, \underline{n} (see, R. J. Charles,

25

Journal of Applied Physics, 29, 1554 (1958)), was approximately 216 as determined through static fatigue tests involving suspending a series of weights from the fiber.

A length of approximately 2 kilometers was placed in one atmosphere of hydrogen at 250 degrees Centigrade. The loss of transmitted light due to the 1.59 µm hydrogen peak was measured as a function of time. After a period, the loss no longer increased. The time to reach half of this loss was considered a measure of the hermeticity of the coating and was approximately 60 hours. This should be compared to a time period of 5 minutes for an identical fiber lacking the hermetic coating.

Example 2

The procedure of Example 1 was followed except that no chlorine was introduced into the reaction chamber. The tensile strength of the fiber was measured by the procedure described in Example 1 and was approximately 500,000 psi.

Example 3

The procedure of Example 1 was followed except the two chamber coating apparatus shown in FIG. 3 was utilized rather than the one chamber apparatus shown in FIG. 2. Nitrogen flows, 33 and 34, were the same as described in Example 1. A nitrogen purge, 31, with a rate of approximately 500 cc/min was utilized to prevent intermixing of reactants from the separate chambers. A reaction mixture containing acetylene, trichloroethylene, and nitrogen in respective mole percentages of 13, 26 and 61 with a total flow rate of 250 cc/min was introduced into the upper chamber at 35. Additionally, a mixture of acetylene and nitrogen was introduced at port, 36, and included a 5 to 95 mole percent composition at a flow rate of 3.5 liters per minute. The fiber was tested as described in Example 1. The static fatigue stress corrosion susceptibility factor, n", was 233, the tensile strength was 602,000 psi, and the half time to saturation of the elevated temperature hydrogen permeation test was approximately that obtained in Example 1.

Claims

1. A process for producing a coated optical fiber comprising the steps of producing a fiber from a heated glass body and contacting said fiber with a gas to induce the formation of a coating on said fiber characterized in that said gas comprises a carbonaceous composition, and said contact oc-

curs when said fiber has a temperature sufficiently high to induce a reaction at the surface of said fiber that results in the formation of an adherent carbon coating comprising a crosslinked carbon network and is sufficiently low to avoid the formation of a diamond-like coating.

10

- The process of claim 1 wherein said gas comprises an alkynyl.
- The process of claim 2 wherein said alkynyl comprises acetylene.
- 4. The process of claim 3 wherein said gas includes a chlorine containing composition.
- 5. The process of claim 4 wherein said chlorine containing composition comprises a gas chosen from the group consisting of trichloroethylene and chlorine.
- 6. The process of claim 5 wherein said coated fiber is subjected to a heat treatment.
- 7. The process of claim 1 wherein said fiber undergoes said contact sequentially, first with a composition comprising a getterer containing gas and said carbonaceous composition and then with a carbonaceous composition in the absence of said getterer.
- 8. The process of claim 7 wherein said carbonaceous composition comprises acetylene.
- 9. The process of claim 7 wherein said getterer comprises a chlorine containing composition.
- 10. The process of claim 9 wherein said chlorine containing composition comprises trichloroethylene.
- 11. The process of claim 1 wherein sald glass body comprises a fiber preform.
- 12. The process of claim 1 wherein said temperature is in the range 700 to 900 degrees C.
- 13. The process of claim 1 wherein said carbon coating is overcoated.
- 14. The process of claim 1 wherein said gas includes an entity chosen from the group consisting of chlorine, bromine, and fluorine.
- 15. The process of claim 1 wherein said fiber contains a getterer for hydrogen or water.
- 16. The product formed by the process of claim 1
- 17. A coated fiber comprising a glass fiber capable of guiding electromagnetic radiation and a coating characterized in that said coating is non-diamond-like, comprises a crosslinked carbon network and has an hermeticity such that the loss measured at 1.24 μ m in 1 atm. of hydrogen and at the use temperature does not increase more than 0.05 dB/km in a time period shorter than 1/140 of the desired use period and such that the static fatigue, $\underline{\bf n}$, value is greater than 70.
- 18. The coated fiber of claim 17 wherein said time period is shorter than 1/45 the desired use period.

55

12

19. A process for producing a coated optical fiber comprising the steps of producing a fiber from a heated glass body and contacting said fiber with a gas to induce formation of an hermetic coating characterized in that a getterer is introduced to remove hydrogen or water trapped by said coating.

ating. I get-

20. The process of claim 19 wherein said getterer is in said fiber.

21. The process of claim 19 wherein sald getterer is introduced in said gas.

22. The process of claim 21 wherein said getterer comprises a chlorine containing compound.

23. The process of claim 22 wherein said chlorine containing compound comprises a gas chosen from the group consisting of trichloroethylene and chlorine.

24. A process for producing a structure comprising a coated optical fiber said process comprising the step of producing a fiber including a core and a cladding from a heated glass body and coating said body characterized in that a getterer for hydrogen or water is introduced into the periphery of said cladding.

25. The process of claim 24 wherein said coating is formed by subjecting said heated fiber to a gas at a temperature sufficient to induce formation of a non-diamond-like crosslinked carbon network.

10

15

20

25

30

35

40

45

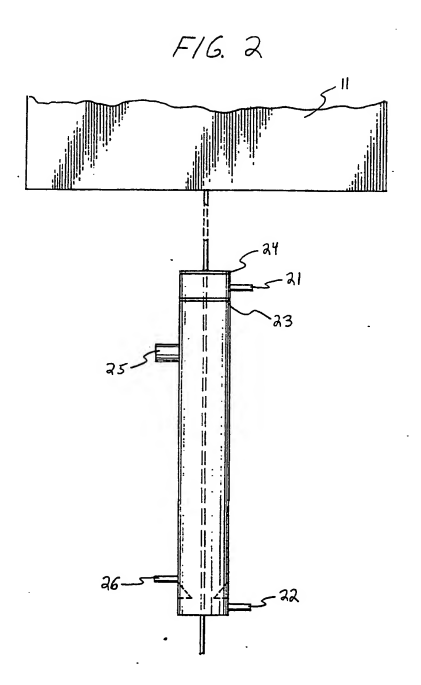
50

EP 0 308 143 A1

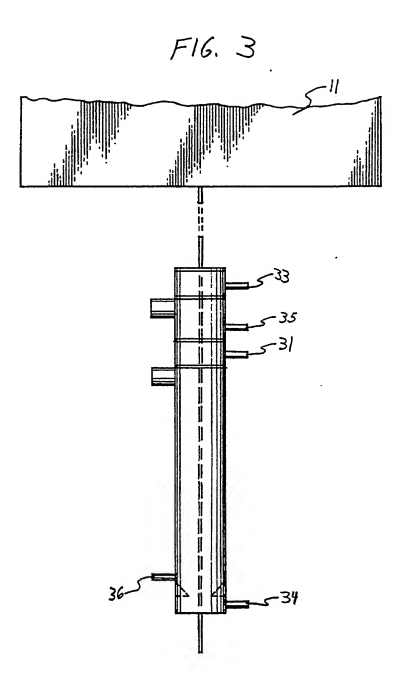
F16. 1

.

EP 0 308 143 A1



EP 0 308 143 A1





EUROPEAN SEARCH REPORT

Application Number

EP 88 30 8335

	DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document with indication, where appropriate, Relevant			elevant	CLASSIFICA	TION OF TH	
Category	of relevant pa			claim	APPLICATIO		
Y	WORLD PATENTS INDEX LATEST DATABASE, accession no. 83-13206K, Derwent Week 06, 1983, Derwent Publicationst Ltd., GB; & JP - A - 57 209 844 (NIPPON T. & T. et al.) 23-12-1982		1,1	11,13 6	C 03 C G 02 B		
Y	GB-A-1 134 133 (QU * page 1, line 64 - claim 1; figures *	ARTZ ET SILICE) page 2, line 10;		3,11- ,16			
Y	FR-A-2 493 302 (LI * claims 1-3 *	GNES T. & T.)		3,11- ,16			
X	WORLD PATENTS INDEX accession no. 83-13 06, 1983, Derwent p GB; & JP - A - 57 2 T.) 23-12-1982	207K, Derwent week	1,	13,16			
X	US-A-4 183 621 (KA * claims 1, 2; colu column 4, lines 5-1	mn 2, lines 17-28;	. 1,	13	TECHNICAL FIELDS SEARCHED (Int. Cl.4)		
A	US-A-4 306 897 (MA * abstract; column		1		C 03 C G 02 B		
A	GB-A-2 156 858 (GE * abstract; claims	C plc) 1, 2, 7-9 *	1,	13			
	The present search report has been present search ERLIN CATEGORY OF CITED DOCUMEN	Date of completion of the search 24-11-1988 T: theory or pr E: earlier pate:	inciple und	erlying the	Examiner PUD J.G. Invention shed on, or		
Y : par doc A : tecl	ilcularly relevant if taken alone ticularly relevant if combined with and ument of the same category nnological background I-written disclosure	L : document ci	ited in the ted for oth	er reasons	77 + 1 100 + 1 100 + 2 + 2 + 2 + 2 + 2 + 2 + 2 + 2 + 2 +		